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Gretchen M. Gallegos Richard A. Brown

SOIL AND SEDIMENT MONITORING

Introduction

The soil and sediment monitoring analysis that Lawrence Livermore National Laboratory performed in 2000 included work in three areas: surface soil in the Livermore Valley and at Site 300, sediment at the Livermore site, and vadose zone soils at the Livermore site.

Soil is weathered material, mainly composed of disintegrated rock and organic material that sustains growing plants. Soil can contain pollutants originally released directly to the ground, to the air, or through liquid effluents. Department of Energy (DOE) guidance for environmental monitoring states that soil should be sampled to determine if there is measurable, long-term buildup of radionuclides in the terrestrial environment and to estimate environmental radionuclide inventories (U.S. DOE 1991). The guidance recommends monitoring for radionuclides specific to a particular operation or facility as well as those that occur naturally. Particulate radionuclides are of major interest in the LLNL soil monitoring program because airborne particulate releases are the most likely pathway for LLNL-induced soil contamination.

Sediments are defined for the purposes of this chapter as finely divided, solid materials that have settled out of a liquid stream or standing water. The accumulation of radioactive materials in sediment could lead to exposure of humans through their ingestion of aquatic species, sediment resuspension into drinking water supplies, inhalation of

dust particles, or as an external radiation source (U.S. DOE 1991). However, the Livermore site and Site 300 do not have habitats for aquatic species that are consumed by people, nor do they have surface drainage that directly feeds drinking water supplies.



Soils in the vadose zone—the region below the land surface where the soil pores are only partially filled with water—are collected in arroyo channels at the Livermore site as part of the Ground Water Protection Management Program (GWPMP). Infiltration of natural runoff through arroyo channels is a significant source of groundwater recharge, accounting for an estimated 42% of resupply for the entire Livermore Valley groundwater basin (Thorpe, et al. 1990). Soils in the shallow vadose zone are collected and analyzed to provide information about possible constituents that may be dissolved as runoff water infiltrates through the arroyo to the groundwater.

Sampling Locations

Since 1971, surface soil sampling near the LLNL Livermore site and Site 300 has been part of a continuing LLNL monitoring program designed to measure any changes in environmental levels of radioactivity and evaluate any increase in radioactivity that might have resulted from LLNL operations. These samples have been analyzed for plutonium and gamma-emitting radionuclides, such as depleted uranium used in some explosive tests at Site 300. The inclusion of other gammaemitting, naturally occurring nuclides (potassium-40 and thorium-232) and the long-lived fission product, cesium-137, provides background information and baseline data on global fallout from historical aboveground nuclear weapons testing. In addition, LLNL analyzes Site 300 soils for beryllium, a potentially toxic metal used at this site. Soils in the Livermore vicinity were analyzed for beryllium from 1991 to 1994. However, analysis for beryllium was discontinued at the Livermore site in 1995, because it was never measured above background values.

Surface soil samples are collected at 19 locations in the Livermore Valley (**Figure 10-1**) and 14 locations at or near Site 300 (**Figure 10-2**). The

locations were selected to represent background concentrations (distant locations unlikely to be affected by LLNL operations) as well as areas where there is the potential to be affected by LLNL operations. Areas with known contaminants, such as the Livermore Water Reclamation Plant (LWRP), are also sampled.

Site 300 soil sampling locations are established around firing tables and other areas of potential soil contamination. PRIM location, east of Site 300, became inaccessible and was removed from the sampling program in 2000 because the site owner discontinued operations.

Sediment samples have been collected from selected arroyos and other drainage areas at and around the Livermore site since 1988; these locations largely coincide with selected storm water sampling locations (see Chapter 7). The sediment sampling location, CDB, was removed in the year 2000; the surface drainage and settling basin at that location had been removed and storm drainage pipe installed, thereby preventing any meaningful sediment sampling. However, four additional samples were collected in the shelf areas of the Drainage Retention Basin, when the basin was partially emptied for the addition of a frog habitat. These samples were available for collection because the water level was maintained during the habitat construction at a level sufficient for sampling.

Sediment sampling locations have not been established at Site 300. The drainage courses at Site 300 are steep, causing flowing water to scour the drainages and preventing the accumulation of sediment. Because of these conditions, sediment sampling at Site 300 is not warranted.

Vadose zone soil sampling has been conducted since 1996. These sampling locations correspond to the same selected storm water sampling locations as the sediment sampling locations (see

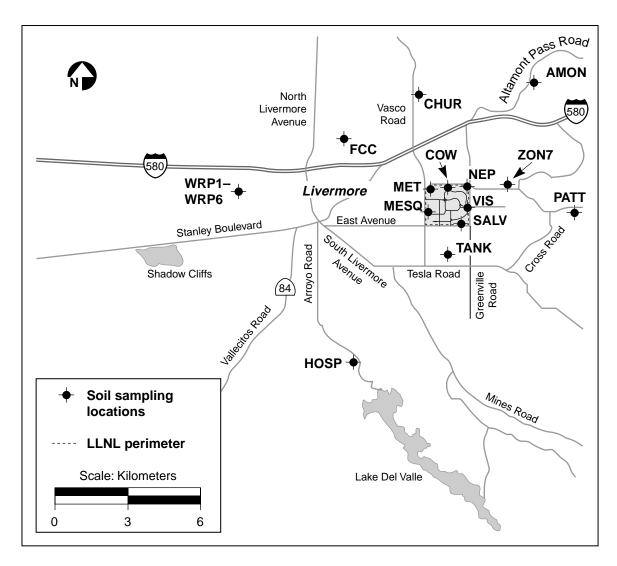


Figure 10-1. Surface soil sampling locations, Livermore Valley, 2000

Figure 10-3). Vadose zone samples were not collected in the Drainage Retention Basin because the liner for the basin prevents migration of materials to the groundwater. The collocation of sampling for these three media facilitates comparison of analytical results. As with sediment samples, vadose zone samples are not collected at Site 300.

Approximately 10% of locations are sampled in duplicate; two identical samples are collected at each location chosen for this sampling. All soil and

sediment sampling locations have permanent location markers for reference.

Methods

Surface soil, sediment, and vadose zone soil sampling is conducted according to written, standardized procedures (Tate et al. 1999). Soil samples are collected from undisturbed areas near permanent location markers. These areas are generally level, free of rocks, and unsheltered by trees or buildings. Surface soil samples are

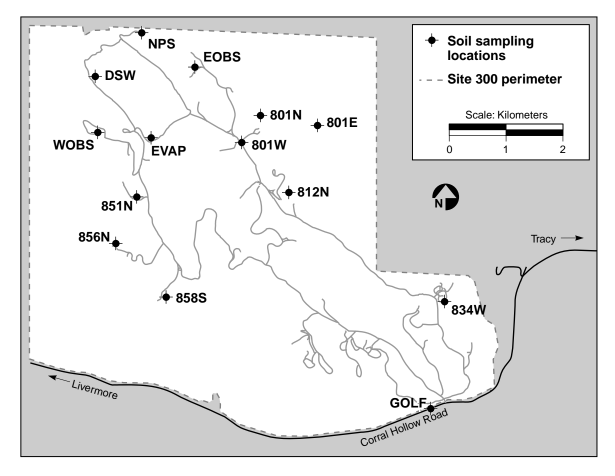


Figure 10-2. Site 300 surface soil sampling locations, 2000

collected from the top 5 cm of soil because aerial deposition is the primary pathway for potential contamination, and resuspension of materials from the surface into the air is the primary exposure pathway to nearby human populations. Sediments are collected annually from drainages at and around the LLNL Livermore site after the cessation of spring runoff. Samples to be analyzed for particulate radionuclides are collected from the top 5 cm of soil. Samples to be analyzed for tritium are collected 5–15 cm deep to obtain sufficient water in the sample for analysis. Vadose zone soil samples are collected at 30-45 cm deep for metals analysis, and at 45-65 cm deep for analysis of soluble volatile organic compounds and for polychlorinated biphenyls (PCBs).

In 2000, surface soil samples in the Livermore Valley were analyzed for plutonium and gammaemitting radionuclides. Samples from Site 300 were analyzed for gamma-emitting radionuclides and beryllium. Analysis of Site 300 soil samples for plutonium was discontinued in 1997 because plutonium has not been used at the site, and sample results have continuously been at background levels since sampling began in 1972. Annual sediment samples collected at the Livermore site were analyzed for plutonium and gamma-emitting radionuclides and tritium. Because the four additional samples from the drainage retention basin were of interest due to the potential for the settling of particulate materials, these samples were analyzed for plutonium and gamma-emitting radionuclides,

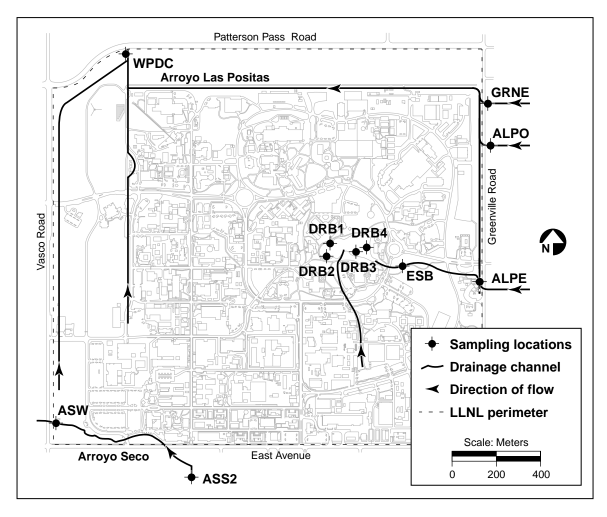


Figure 10-3. Sediment and vadose zone sampling locations on or near the Livermore site, 2000

but not for tritium. Vadose zone samples were analyzed for total and soluble metals and for soluble volatile organic compounds.

Prior to radiochemical analysis, surface soil and sediment samples are dried, ground, sieved, and homogenized. The samples are analyzed by LLNL's Chemistry and Materials Science Environmental Services (CES) laboratory. The plutonium content of a 100-g sample aliquot is determined by alpha spectroscopy. Other sample aliquots (300-g) are analyzed for more than 150 radionuclides by gamma spectroscopy using a high-purity germanium (HPGe) detector (Hall and Edwards 1994a,

b, and c). The 10-g subsamples for beryllium analyses are sent to a contract analytical laboratory and are analyzed by graphite-furnace atomic absorption spectroscopy. For sediment samples collected for tritium analyses, CES uses freezedrying techniques to recover water from the samples and determines the tritium content of the water by liquid-scintillation counting.

Vadose zone soil samples are analyzed by a contract laboratory. The analytical methods include the toxicity characteristic leaching procedure (TCLP) followed by EPA Method 8240 for volatile organic compounds, and total metals by EPA Methods

200.7, 245.2, 7471A, and 6010B. The procedure for determining soluble metals includes the California Waste Extraction Test, followed by the same analytical methods for metals applied to the leachates. In 2000, a vadose zone soil sample from location ESB (**Figure 10-3**) was also analyzed for PCBs by EPA Method 8082. Chain-of-custody procedures are followed throughout the sampling, delivery, and analytical processes.

Livermore Valley Surface Soil Results

Table 10-1 presents summary data on the concentrations of plutonium-239+240, plutonium-238, americium-241, cesium-137, potassium-40, thorium-232, uranium-235, and uranium-238 in surface soils from the Livermore Valley sampling locations. Complete data for 2000 soil and sediment sampling are presented in Table 10-1 of the Data Supplement.

Table 10-1. Summary of surface soil and sediment radioanalytical data, 2000

Analyte and location	Detection frequency ^(a)	Median	IQR ^(b)	Maximum
Plutonium-238 (µBq/dry g)				
Livermore Valley soils	12/13	2.70	3.65	19.5
LWRP ^(c) soils	6/6	128	173	377
Livermore site sediments	6/10	6.53	11.8	174
Plutonium-239+240 (µBq/dry g)				
Livermore Valley soils	13/13	61.1	60.3	352
LWRP soils	6/6	2410	3370	6440
Livermore site sediments	10/10	78.9	105	1740
Cesium-137 (mBq/dry g)				
Livermore Valley soils	13/13	1.39	0.960	7.1
LWRP soils	6/6	1.04	1.89	4.07
Livermore site sediments	10/10	0.184	0.330	1.11
Site 300 soils	13/14	1.56	1.60	5.00
Potassium-40 (Bq/dry g)				
Livermore Valley soils	13/13	0.470	0.129	0.551
LWRP soils	6/6	0.358	0.045	0.411
Livermore site sediments	10/10	0.391	0.270	0.470
Site 300 soils	14/14	0.414	0.107	0.551
Thorium-232 (μg/dry g) ^(d)				
Livermore Valley soils	13/13	6.73	1.12	8.68
LWRP soils	6/6	6.32	0.06	6.45
Livermore site sediments	10/10	3.16	1.50	6.81
Site 300 soils	14/14	8.08	1.21	11.1



Table 10-1. Summary of surface soil and sediment radioanalytical data, 2000 (continued)

Analyte and location	Detection frequency ^(a)	Median	IQR ^(b)	Maximum
Uranium-235 (µg/dry g) ^(e)				
Livermore Valley soils	13/13	0.0190	0.0057	0.0272
LWRP soils	6/6	0.0190	0.0030	0.0237
Livermore site sediments	10/10	0.0106	0.0039	0.0175
Site 300 soils	14/14	0.0301	0.0117	0.0690
Uranium-238 (µg/dry g) ^(f)				
Livermore Valley soils	13/13	1.78	0.750	2.87
LWRP soils	6/6	1.94	0.435	2.74
Livermore site sediments	10/10	1.02	0.464	1.57
Site 300 soils	14/14	1.90	1.41	29.3
Tritium (Bq/L extracted water) ^(g)				
Livermore site sediments	4/6	4.53	1.84	14.1
Americum-241 (mBq/dry g) ^(h)				
LWRP soils	2/6	1.00	(i)	3.17
Beryllium (mg/kg) ^(j)				
Site 300 soils	9/14	0.36	(i)	1.10

- a The detection frequency is the fraction of samples having a measured value above the detection limit.
- b IQR = Interquartile range; the difference between the top of the third and the top of the first quartiles of the data
- c LWRP = Livermore Water Reclamation Plant
- d Thorium-232 activities in Bq/dry g can be determined by dividing the weight in μg/dry g by 247.3, and pCi/dry g can be determined by dividing by 9.15.
- e Uranium-235 activities in Bq/dry g can be determined by dividing the weight in μg/dry g by 12.5, and pCi/dry g can be determined by dividing by 0.463.
- f Uranium-238 activities in Bq/dry g can be determined by dividing the weight in μg/dry g by 80.3, and pCi/dry g can be determined by dividing by 2.97.
- g Only sediment samples are analyzed for tritium. The samples collected within the Drainage Retention Basin were not analyzed for tritium because only particulate materials were of interest in these samples.
- h Americium-241 is detected only in LWRP soil samples.
- i IQR is not calculated because of high incidence of reported values below detection limits.
- j Only Site 300 samples are analyzed for beryllium.

The concentrations and distributions of all observed radionuclides in soil for 2000 are within the ranges reported in previous years and generally reflect worldwide fallout and naturally occurring concentrations. The ratio of uranium-235 to uranium-238 generally reflects the natural ratio of 0.7%; however, there is significant uncertainty in the uranium-235/uranium-238 ratio because of

the difficulty in measuring small quantities of uranium-238 by gamma spectroscopy.

Plutonium has, in the past, been detected at levels above background at VIS, a perimeter sampling location near the east boundary of the Livermore site. Since 1980, soil samples at this location have generally shown plutonium-239+240 values higher

than background. However, in 2000, the measured plutonium-239+240 value for VIS was within the range of background, 352 μBq/dry g $(9.51 \times 10^{-3} \,\mathrm{pCi/dry}\,\mathrm{g})$. The slightly higher values at and near the Livermore site have been attributed to historic operations, including the operation of solar evaporators for plutoniumcontaining liquid waste in the southeast quadrant (Silver et al. 1974). LLNL no longer operates the solar evaporators or engages in any other open-air treatment of plutonium-containing waste. Nonetheless, plutonium-239+240 from historic operations can be carried off site by resuspension of soil by wind. Similarly, elevated levels of plutonium-239+240 (resulting from an estimated 1.2×10^9 Bq [32 mCi] plutonium release to the sanitary sewer in 1967 and earlier releases) were first observed in soils near LWRP during the early 1970s, and were again detected at LWRP sampling locations. As in 1997 through 1999, americium-241 was detected in LWRP samples; it is most likely caused by the natural decay of the trace concentrations of plutonium-241 that were

Historical plots of median plutonium-239+240 concentrations in soil in the Livermore Valley upwind and downwind of the center of the LLNL Livermore site, at Site 300, and at LWRP are shown in **Figure 10-4**. Livermore Valley upwind and Site 300 concentrations have remained relatively constant since monitoring began and generally are indicative of worldwide fallout. Greater variation can be noted in the downwind concentration data, which in 2000 included sampling locations VIS, PATT, NEP, COW, AMON, and ZON7, compared with the upwind and historic Site 300 data. The concentrations of plutonium at the downwind locations reflect resuspension of low-level plutonium contamination from soils in the southeast quadrant of the Livermore site. Greater variability in plutonium-239+240 is seen in samples from LWRP.

present in the release.

Because the plutonium-239+240 is likely to be present in discrete particles, the random presence or absence of the particles dominates the measured plutonium-239+240 in any given sample.

Livermore Site Sediment Results

Table 10-1 presents summary data on radionuclides detected in the sediment samples; a complete presentation of 2000 sediment data is found in Table 10-1 of the Data Supplement. The levels of plutonium-239+240 were generally at background concentrations, reflective of worldwide fallout. Sampling location ESB (see Figure 10-3) shows a moderately higher value for plutonium than values at other locations. The value may be attributed to historic actions because this location is in a drainage area for the southeast quadrant at LLNL. Most other radionuclides were detected at levels similar to those reported from 1988 through 1999: cesium-137 (a fission product) was found at worldwide background concentrations; potassium-40, thorium-232, uranium-235, and uranium-238 (naturally occurring radionuclides) were detected at background concentrations. Tritium concentrations were within the range of previous data. Tritium in sediments will continue to be evaluated as long as the measured values remain above the detection limits of the liquid scintillation analytical method.

Livermore Site Vadose Zone Soil Results

Analytical results for vadose zone soil samples are compared with soil reuse standards developed by LLNL and approved by the San Francisco Bay Regional Water Quality Control Board (SFBRWQCB) (Folks 1997, Marshack 2000). Metals background concentrations are based on naturally occurring levels in the soil, considering first the results for total metals and then the soluble metals test. There are no background levels for organic compounds or tritium. Soils containing

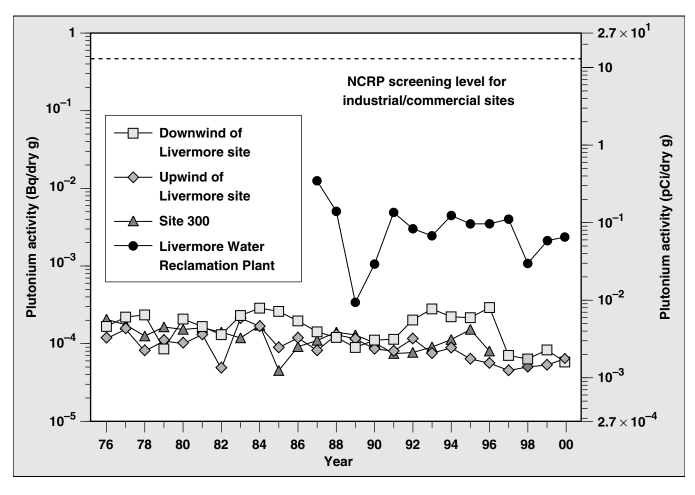


Figure 10-4. Median plutonium-239+240 activities in surface soils, 1976–2000. Upwind and downwind designations are relative to the center of the Livermore site.

materials at levels above background still may not adversely affect the groundwater. If there are any detected organic compounds or tritium, the designated level methodology (DLM) (i.e., application of a simple attenuation factor and specific water quality objectives) is used to determine the soluble levels of contaminants that would not adversely impact groundwater beyond its beneficial uses. (Background and DLM de minimis values are presented in Tables 10-3 and 10-4 in the Data Supplement.)

All analytical results for soluble VOCs were below detection limits, except for chloroform, a common laboratory contaminant. Chloroform was also detected in the laboratory's method blank samples. All total metals concentrations were within site background. See Tables 10-5 to 10-7 in the Data Supplement for analytical results for VOCs and metals. A PCB, Arochlor 1260, was detected at 14 mg/kg at location ESB. The presence of PCBs suggests that this sample represents residual low-level contamination from the 1984 excavation of the former East Traffic Circle landfill (see Chapter 9). Further confirmatory investigations are underway. (The analytical results for approximately the same location sampled in 2001 was 2.1 mg/kg.) The detected concentrations are below the federal and state hazardous waste limits. Tritium results from the sediment sampling were evaluated by the DLM

method and were all below de minimis levels (see the Data Supplement, Table 10-1).

Site 300 Results

Table 10-1 presents summary data on the concentrations of cesium-137, potassium-40, thorium-232, uranium-235, uranium-238, and beryllium in soil from the Site 300 sampling locations; a complete presentation of 2000 soils data for

Site 300 is found in Table 10-2 of the Data Supplement. The concentrations and the distributions of all observed radionuclides in Site 300 soil for 2000 lie within the ranges reported in all years since monitoring began. The ratio of uranium-235 to uranium-238 generally reflects the natural ratio of 0.7%. Historical trends of uranium-238 concentrations from both the Livermore Valley and Site 300 are shown in **Figure 10-5**. Median values have remained relatively constant for both places.

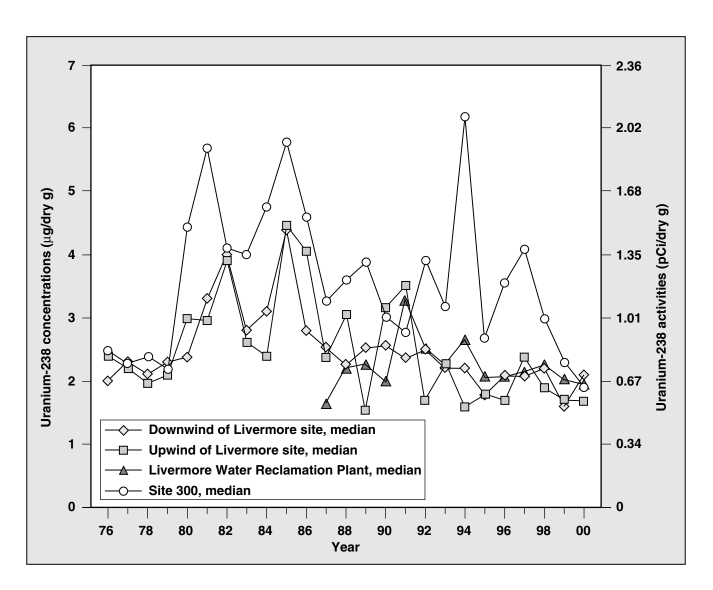


Figure 10-5. Uranium-238 concentrations in surface soils, 1976–2000

The highest values at Site 300 result from the use of depleted uranium in explosive experiments.

Environmental Impact

This section discusses the environmental impacts of operations at the LLNL Livermore site and Site 300 inferred from soil, sediment, and vadose zone soil monitoring.

Livermore Site

Routine surface soil, sediment, and vadose zone soil sample analyses indicate that the impact of LLNL operations on these media in 2000 has not changed from previous years and remains insignificant. Most analytes of interest or concern were detected at

background concentrations or in trace amounts, or could not be measured above detection limits.

The highest value of 6.4 mBq/dry g (0.17 pCi/dry g) for plutonium-239+240 measured at LWRP is 1.3% of the National Council on Radiation Protection (NCRP) recommended screening limit of 470 mBq/g (12.7 pCi/g) for property used for commercial purposes (NCRP 1999). Statistical analysis shows no general increase or decrease in plutonium-239+240 values with time.

Over the years, LLNL has frequently investigated the presence of radionuclides in local soils. Several of the studies are listed in **Table 10-2**. LLNL sampling of surface soil, sediment, and vadose zone soil will continue on an annual basis.

Table 10-2. Special soil studies

Year	Subject	Reference
1971–1972	Radionuclides in Livermore Valley soil	Gudiksen et al. 1972; Gudiksen et al. 1973
1973	Radionuclides in San Joaquin Valley soil	Silver et al. 1974
1974	Soil study of southeast quadrant of Livermore site	Silver et al. 1975
1977	Sediments from LLNL to the San Francisco Bay	Silver et al. 1978
1980	Plutonium in soils downwind of the Livermore site	Toy et al. 1981
1990	195 samples taken in southeast quadrant for study	Gallegos et al. 1992
1991	Drainage channels and storm drains studied	Gallegos 1991
1993	EPA studies southeast quadrant	Gallegos et al. 1994
1993	Historic data reviewed	Gallegos 1993
1995	LLNL, EPA, and DHS sample soils at Big Trees Park	MacQueen 1995
1999	Summary of results of 1998 sampling at Big Trees Park	Gallegos et al. 1999
2000	Health Consultation, Lawrence Livermore National Laboratory, Big Trees Park 1998 Sampling	Agency for Toxic Substances Disease Registry 2000

Site 300

The concentrations of radionuclides and beryllium observed in soil samples collected at Site 300 are within the range of previous data and are generally representative of background or naturally occurring levels. The uranium-235/uranium-238 ratios that are indicative of depleted uranium occur near active and inactive firing tables at Buildings 801 and 812. There they represent a small fraction of the firing table operations that disperse depleted uranium. The uranium-238 concentrations are below the NCRP recommended screening level for commercial sites of 313 μ g/g (3.9 Bq/g or 105 pCi/g). Historically, some measured concentrations of uranium-238 near Building 812 have been greater than the screening level. The investigation and characterization planned for the area surrounding Building 812 will clarify the nature and extent of the contamination in the area.